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(11) EP 0 846 654 A1

(12)

## **EUROPEAN PATENT APPLICATION**

(43) Date of publication: 10.06.1998 Bulletin 1998/24

(51) Int Cl.6: C01B 15/013

(21) Application number: 97402937.3

(22) Date of filing: 04.12.1997

(84) Designated Contracting States:

AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC

NL PT SE

Designated Extension States:

AL LT LV MK RO SI

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## (54) On-site manufacture of ultra-high-purity hydrogen peroxide

(57) Provided is a novel on-site system and method for providing ultra-high-purity reagents including  $H_2O_2$  to a point of use. The subsystem includes: (a) a tank connected to receive aqueous  $H_2O_2$  and to provide a flow of  $H_2O_2$  therefrom; (b) an anionic exchange bed and a cationic exchange bed connected to receive the flow of  $H_2O_2$  from the tank, and to produce therefrom a purified  $H_2O_2$  flow with a reduced level of ionic contaminants. The cationic exchange bed is preconditioned

with acid and the anionic bed is preconditioned with bicarbonate ions; (c) a filter downstream of the anionic exchange bed and the cationic exchange bed; and (d) piping which routes the aqueous  $H_2O_2$  from the filter to the point of use. The inventive system and method have particular applicability in semiconductor fabrication, for supplying ultra-high-purity hydrogen peroxide and hydrogen peroxide-containing reagents to a point of use, such as a wafer cleaning/etching stations.

#### Description

#### CROSS REFERENCE TO RELATED APPLICATIONS

This application is a continuation-in-part of copending application Serial No. 08/675,630, filed on July 1, 1996, which is herein incorporated by reference.

#### BACKGROUND OF THE INVENTION

#### 1. Field of the Invention

The present invention relates to various on-site subsystems for providing ultra-high-purity reagents comprising hydrogen peroxide to a point of use. The present invention also relates to methods for the on-site provision of ultra-high-purity reagents comprising hydrogen peroxide to a point of use. The invention has particular applicability in semiconductor fabrication, for providing ultra-high-purity materials to a semiconductor manufacturing operation.

#### 2. Description of the Related Art

#### a. Contamination Control

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Contamination is generally an overwhelmingly important concern in integrated circuit (IC) manufacturing. A large fraction of the steps used in modern integrated circuit manufacturing are cleanup steps of one kind or another. Such cleanup steps are used, for example, to remove organic contaminants, metallic contaminants, photoresist (or inorganic residues thereof), byproducts of etching, native oxides, etc.

As of 1995, the cost of a new front end (IC wafer fabrication facility) was typically more than one billion dollars (\$1,000,000,000). A large fraction of the cost for such facilities is directed to measures for particulate control, cleanup, and contamination control.

One important and basic source of contamination in semiconductor fabrication is impurities in the process chemicals. Since the cleanup steps are performed so frequently in and are so critical to IC fabrication, contamination due to cleanup chemistry is very undesirable.

#### b. Wet versus Dry Processing

One of the long-running technological shifts in semiconductor processing has been the changes (and attempted changes) between dry and wet processing. In dry processing, only gaseous or plasma-phase reactants come in contact with the wafer or wafers being treated. In wet processing, a variety of liquid reagents are used for a multitude of purposes, such as the etching of silicon dioxide, silicon nitride and silicon, and the removal of native oxide layers, organic materials, trace organic or inorganic contaminants and metals.

While plasma etching has many attractive capabilities, it is not adequate for use in cleanup processes. There is simply no available chemistry with plasma etching to remove some of the most undesirable impurities, such as gold. Thus, wet cleanup processes are essential to modern semiconductor processing, and are likely to remain so for the foreseeable future.

Plasma etching is performed using a photoresist mask in place, and is not immediately followed by high-temperature processes. After plasma etching, the resist is stripped from the wafer surface using, for example, an  $O_2$  plasma treatment. Cleanup of the resist stripped wafer(s) is then necessary.

The materials which the cleanup process should remove include, for example, photoresist re'sidues (organic polymers), sodium, alkaline earth metals (e.g., calcium, magnesium) and heavy metals (e.g., gold). Many of these contaminants do not form volatile halides. As a result, plasma etching will not remove such contaminants from the wafer surface. Hence, cleanup processes using wet chemistries are required.

Because any dangerous contaminants stemming from the plasma etching process are removed prior to high-temperature processing steps by wet chemical treatment, the purities of plasma etching process chemicals (i.e., liquified or compressed gases) are not as critical as those of the liquid chemicals used in cleanup processes. This difference is due to the impingement rate of the liquid chemical at the semiconductor surface typically being one million times greater than that of the plasma species in plasma etching. Moreover, since the liquid cleanup steps are directly followed by high-temperature processes, contaminants on the wafer surface tend to be driven (i.e., diffused) into the wafer.

Wet processing has a major drawback insofar as ionic contamination is concerned. Integrated circuit devices generally use only a few dopant species (e.g., boron, arsenic, phosphorus, and antimony) to form the requisite p-type and n-type doped regions of the device. However, many other species act as electrically active dopants, and are highly

undesirable contaminants. These contaminants can have deleterious effects on the IC devices, such as increased junction leakage at concentrations well below 1013cm-3.

Moreover, some less desirable contaminants segregate into the silicon substrate. This occurs when silicon is in contact with an aqueous solution, and the equilibrium concentration of the contaminants is higher in the silicon than in the solution. Moreover, some less desirable contaminants have very high diffusion coefficients. Consequently, introduction of such contaminants into any part of the silicon wafer may result in diffusion of the contaminants throughout the water including junction locations where leakage may result.

Thus liquid solutions for treating semiconductor wafers should have extremely low levels of metal ions. Preferably, the concentration of all metals combined should be less than 300 ppt (parts per trillion), and less than 10 ppt for any single metal. Even lower concentrations are desirable. Contamination by anions and cations should also be controlled. Some anions may have adverse effects, such as complexed metal ions which reduce to mobile metal atoms or ions in the silicon lattice.

Front end facilities typically include on-site purification systems for preparation of high-purity water (i.e., "deionized" or "DI" water). However, it is more difficult to obtain liquid process chemicals in the purities required.

#### c Purity in Semiconductor Manufacturing

Uncetected contamination of chemicals increases the probability for costly damage to a large quantity of wafers. The extreme purity levels required by semiconductor manufacturing are rare and unique among industrial processes. With such extreme purity requirements, handling of chemicals is undesirable (though of course it cannot be entirely avoided). Exposure of ultrapure chemicals to air (particularly in an environment where workers are also present) should be minimized. Such exposure risks the introduction of particulates into the chemicals, which can result in the contamination of those chemicals. Furthermore, shipment of ultrapure chemicals in closed containers is not ideal, since such containers increase the risk of contaminants being generated at the manufacturer's or at the user's site.

Since many corrosive and/or toxic chemicals are used in semiconductor processing, the reagent supply locations are commonly separated from the locations where front-end workers are present. Construction and maintenance of piping for ultra-high-purity (UHP) gases and liquids are well-understood in the semiconductor industry, and most gases and liquids can be transported to wafer fabrication stations from anywhere in the same building (or in the same site).

#### d Hydrogen Peroxide

Hydrogen peroxide  $(H_2O_2)$  is an important process chemical in the semiconductor manufacturing industry. It is commonly used in cleanup solutions. For example, the widely used "piranha" cleanup solution typically uses  $H_2O_2 + H_2SO_4$  in proportions of 30 70, and the widely used "RCA" cleanup is a three-stage cleanup which uses hydrogen peroxide in two of the stages

Thus, ultra-high-purity (UHP) aqueous hydrogen peroxide is a staple to integrated circuit (IC) manufacturing. Hydrogen peroxide is not an easy chemical to purify, however, since its decomposition is exothermic and temperature sensitive as well as being catalyzed by various metals and contaminants. Moreover, it is known that  $H_2O_2$  is a powerful oxidant. Substantial work has been done in this area. For example, it has been reported in the literature that an anionic exchange resin for purifying hydrogen peroxide should preferably be loaded with bicarbonate (HCO<sub>3</sub>-) ions, since other commonly used anions (e.g., OH- and C $\ell$ -) will catalyze the decomposition of  $H_2O_2$  under some circumstances.

Elimination of organic acidic components contained in  $H_2O_2$  is also described in the literature. See, e.g., French Patent Application No. 1,539.843 (using non-functionalized resins and neutralized base acid); U.S. Patent No. 3,294.488 (basic resin [HCO<sub>3</sub>] + CO<sub>2</sub>); Japanese Patent No. 6,725,845 (1967) (non-functionalized resins); U.S. Patent No. 297.404 (basic resin [HCO<sub>3</sub>] and CO); U.S. Patent No. 3,305,314 (basic resin [HCO<sub>3</sub>] and CO<sub>2</sub>/3\*); and U.S. Patent No. 4,792,403 (halogenated resin).

H<sub>2</sub>O<sub>2</sub> purification on cationic and anionic resins (tertiary, quaternary) is described in, for example, French Patent Application No. 10,431 (1953) (use of Sulfonic resins); Polish Patent No. 50,982 (1961) (cationic + anionic resins); Polish Patent No. 55,378 (1968); Spanish Patent No. 328,719 (1961) (sulfonic resins, acrylic, strong base and acid [gel type]); U.S. Patent No. 3,297,404 (1967) (use of mixed resins cationic and anionic [HCO<sub>3</sub>]; and U.S. Patent No. 4,999,179 (sulfonic resin + anionic resin [HCO<sub>3</sub>], CO2/3 + brominated). Various configurations are described, for example, in French Patent No. 2,677,010 (1992) (strong cationic resin + medium strength anionic resin of the gel type + non-functionalized resin); French Patent No. 2,677,011 (1992) (medium strength anionic resin); and PCT Application No. 92/06918 (1992) (cationic and anionic resins, description of fluidized bed technology).

 $H_2O_2$  purification on sulfonic and pyridinic resins is described in, for example, Swedish Patent No. 1,643,452 (1991) (cationic resin + 2,5 methyl base resins - pyridinic vinyl [HCO<sub>3</sub><sup>-</sup>]); and Japanese Patent No. 62,187,103 (1966) (cationic resins + pyridilic anionic structure).

H<sub>2</sub>O<sub>2</sub> purification on resins and chelatants is described in, for example, French Patent No. 2,624,500 (1988) (adding

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carboxylic or phosphonic chelatant to the basic resin); German Patent No. 3,822,248 (1990) (EDTA added to basic resin); European Patent No. 502,466 (1992) (Chelatant added to  $H_2O_2$  and passing into non-functionalized resin); U. S. Patent No. 5,200,166 (addition of stabilizing acid into  $H_2O_2$  and reacting with basic resin [HCO<sub>3</sub><sup>-</sup>, CO<sub>2</sub>/<sub>3</sub><sup>-</sup>]); European Patent No. 626,342 (1994) (chelatant resin with phosphates <0.1 ppm A<sup>-</sup> or A<sup>-</sup>/C<sup>+</sup> + chelatant Aℓ and Fe).

Several patents have also reported success in achieving purities below 1 part per billion (ppb). See, e.g., French Patent No. 624,500 (using resins and complexing agents); WO90/11967 (using SnO<sub>2</sub> + ultrafiltration); and French Patent Application No. 3,045,504 (using Silica treatment). All of the above-cited documents are hereby incorporated by reference.

#### e. Objects and Advantages of the Invention

To meet the requirements of the semiconductor processing industry and to overcome the disadvantages of the related art, it is an object of the present invention to provide novel on-site subsystems for providing ultra-high-purity reagents comprising  $H_2O_2$  to a point of use, in which the hydrogen peroxide can be piped directly to the point of use. The system is very compact, and can be located in the same building as the point of use (or in an adjacent building), such that chemical handling can be avoided. As a result, low impurity levels on a semiconductor wafer surface can be achieved. Ultimately, improvements in device characteristics and increased product yield can be realized.

It is a further object of the invention to provide methods for the on-site provision of ultra-high-purity reagents comprising  $H_2O_2$  to a point of use.

It is, additionally, an object of the present invention to provide an on-site subsystem and method, in a semiconductor device fabrication facility, for providing ultra-high-purity reagents comprising  $H_2O_2$  to a semiconductor manufacturing operation.

Other objects and aspects of the present invention will become apparent to one of ordinary skill in the art upon review of the specification, drawings and claims appended hereto.

#### SUMMARY OF THE INVENTION

The foregoing objectives are met by the systems and methods of the present invention. According to a first aspect of the present invention, a novel on-site system for providing ultra-high-purity reagents comprising  $H_2O_2$  to a point of use is provided. The subsystem includes: (a) a tank connected to receive aqueous  $H_2O_2$  and to provide a flow of  $H_2O_2$  therefrom; (b) an anionic exchange bed and a cationic exchange bed connected to receive the flow of  $H_2O_2$  from the tank, and to produce therefrom a purified  $H_2O_2$  flow with a reduced level of ionic contaminants. The cationic exchange bed is preconditioned with acid and the anionic bed is preconditioned with bicarbonate ions; (c) a filter downstream of the anionic exchange bed and the cationic exchange bed; and (d) piping which routes the aqueous  $H_2O_2$  from the filter to the point of use.

According to a second aspect of the invention, a novel on-site subsystem for providing ultra-high-purity reagents comprising H<sub>2</sub>O<sub>2</sub> to a point of use is provided, in which a mixed anionic/cationic exchange bed is utilized.

According to a third aspect of the invention, an on-site subsystem for providing ultra-high-purity reagents comprising  $H_2O_2$  to a point of use is provided. This subsystem includes the following features: (a) a tank connected to receive aqueous  $H_2O_2$  and to provide a flow of  $H_2O_2$  therefrom; (b) an anionic exchange bed and a cationic exchange bed connected to receive the flow of  $H_2O_2$  from the tank, and to produce therefrom a purified  $H_2O_2$  flow with a reduced level of ionic contaminants; (c) an ionic purifier system, connected to pass a gaseous reagent precursor through an area of gas/liquid contact to produce an ultrapure gaseous reagent; (d) generation and mixing subsystems, connected to combine the ultrapure gaseous reagent with deionized water and with the purified  $H_2O_2$  flow to produce an ultrapure cleanup solution; and (e) piping which routes the ultrapure cleanup solution to the point of use.

According to a fourth aspect of the invention, a novel method for the on-site provision of ultra-high-purity reagents comprising  $H_2O_2$  to a point of use is provided. The method includes the steps of: (a) providing a flow of aqueous  $H_2O_2$  from a tank which is located at the same site as the point of use; (b) passing the flow of  $H_2O_2$  through an anionic exchange bed and a cationic exchange bed, to produce a purified  $H_2O_2$  flow with a reduced level of ionic contaminants, wherein the cationic exchange bed is preconditioned with acid and the anionic bed is preconditioned with bicarbonate ions; (c) filtering the purified flow, to produce a flow of ultra-purity aqueous  $H_2O_2$  solution; and (d) routing the flow of ultra-purity aqueous  $H_2O_2$  solution through piping to the point of use.

According to a fifth aspect of the invention, a method for the on-site provision of ultra-high-purity reagents comprising  $H_2O_2$  to a point of use is provided, in which a flow of  $H_2O_2$  passes through a mixed anionic/cationic exchange bed

According to a sixth aspect of the invention, a novel method for the on-site provision of ultra-high-purity reagents comprising  $H_2O_2$  to a point of use is provided. The method includes the steps of: (a) providing a flow of aqueous  $H_2O_2$  from a tank which is located at the same site as the point of use; (b) passing the flow of  $H_2O_2$  through an anionic

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exchange bed and a cationic exchange bed, to produce a purified H2O2 flow with a reduced level of ionic contaminants; wherein the cationic exchange bed is preconditioned with acid and the anionic bed is preconditioned with bicarbonate ions; (c) passing a gaseous reagent precursor through an area of gas/liquid contact in an ionic purifier system to produce an ultrapure gaseous reagent; (d) combining the ultrapure gaseous reagent with deionized water and with the purified H2O2 flow to produce an ultrapure cleanup solution; and (e) routing the flow of ultra-purity cleanup solution to the point of use.

#### BRIEF DESCRIPTION OF THE DRAWING

The objects and advantages of the invention will become apparent from the following detailed description of the preferred embodiments thereof in connection with the accompanying drawings, in which:

FIG. 1 shows an on-site system for purification of aqueous hydrogen peroxide;

FIG. 2 illustrates a process flow for generating an RCA cleanup solution on-site at a wafer fabrication facility; and FIG. 3 is a block diagram of semiconductor cleanup stations in a wafer fabrication facility, in which the purification system of FIG. 1 may be incorporated.

#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS OF THE INVENTION

A first embodiment of the invention will now be described with reference to FIG. 1, which illustrates an on-site 20 system and process flow used for hydrogen peroxide ultrapurification. System 1 can be used by itself simply for the preparation of UHP H<sub>2</sub>O<sub>2</sub>. Alternatively, the system can be used to provide chemicals directly or indirectly to semiconductor process tools.

In accordance with the present invention, incoming  $H_2O_2$  can be purified to impurity levels down to the ppb level. Typical H<sub>2</sub>O<sub>2</sub> target impurities according to the invention are as follows:

- cation concentration < 1.0 ppb;
- anion concentration < 20 ppb; and
- total organic contaminants < 20 ppm.

Aqueous  $H_2O_2$  supply 2 is provided in the system. The concentration of the  $H_2O_2$  supply is preferably high-purity (a feed concentration of 30%  $H_2O_2$  is shown in FIG. 1 for illustration). In accordance with the invention, incoming aqueous  $H_2O_2$  supply 2 is further purified in on-site purification units before being made available for combination with other reagents and subsequent feed to process tools. The on-site purification units preferably include an anionic exchange bed 3 and a cationic exchange bed 4, together with one or more particulate filters 5. Suitable particulate filters are well known to persons skilled in the art.

The containers and piping used in the ultrapurification system are preferably selected from among materials which are inert and non-catalytic to  $H_2O_2$ . Inert fluoropolymers are preferred, since most metals catalyze  $H_2O_2$  decomposition at least to some extent.

In a preferred embodiment of the on-site ultrapurification system, an anionic exchange column 3 is used in combination with a cationic exchange column 4. However, other conventional techniques for sub-ppb polishing can alter-

The ultrapurification system further preferably has a filtration stage 5 disposed downstream from the anion and cation exchange resin columns 3 and 4. Filter stage 5 allows for the removal of any particulates which may have been introduced by columns 4 and 5.

The anionic exchange column is preferably preconditioned by initially loading the column with bicarbonate ions. Bicarbonate preconditioning is described, for example, in U.S. Patent Nos. 3,294,488 and 3,305,314, which are herein incorporated by reference. The bicarbonate preconditioning step is preferably achieved by use of a concentrated NH<sub>4</sub>HCO<sub>3</sub> solution. Possible alternatives to the use of NH<sub>4</sub>HCO<sub>3</sub> include, for example, use of an alkali bicarbonate, which requires removal of the alkali metal ions, or use of CO2, which is inefficient due to the low solubility of CO2.

In a preferred embodiment, the anionic resin is DOWEX MONOSPHERE A550 UPN (polystyrene - DVB gel, quaternary ammonium, 1.0 eq OH-/l) nuclear grade, from Dow. Other anionic resins, however, can be used, such as AMBERLITE IRA 958, from Rohm and Haas (polyacrylic - DVB macroporous, quaternary ammonium 0.8 eq. Ct/ℓ). The cationic exchange column 4 is preferably preconditioned with an acid. This can be done, for example, by washing the cationic exchange column with H<sub>2</sub>SO<sub>4</sub> (e.g., a 10% solution). According to a preferred embodiment, the cationic resin is DOWEX MONOSPHERE C650 UPN (polystyrene - DVB gel, sulfonic, 1.9 eq. H<sup>-</sup>/ℓ), from Dow. Additional suitable cationic resins include, for example, IRA 963 from Rohm and Haas.

While the ultrapurification system illustrated in FIG. 1 depicts a single anionic exchange column and a single

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cationic exchange column, the system preferably has a plurality of anionic and/or cationic exchange columns. In such a case, the plural anionic exchange columns can be placed in series with or in parallel to each other. The plural cationic exchange columns can be configured in a similar manner. In a preferred embodiment, there are two anionic and two cationic exchange columns in series, with the two anionic columns preceding the cationic exchange columns. Other configurations of plural exchange columns will be apparent to those skilled in the art.

Additionally or alternatively to the use of separate anionic and cationic exchange columns, one or more anionic/ cationic mixed beds can be used. When a mixed bed is used in addition to the anionic and cationic exchange columns, the mixed bed is preferably disposed downstream of the anionic and cationic exchange columns.

Suitable mixed bed resins are known in the art, and include, for example, 50/50 DOWEX MONOSPHERE A550 and C650. In the event a mixed bed is used, the resins cannot be preconditioned in situ but must be conditioned prior to packing into the bed.

The following example illustrates the effects of different types of resins or elemental impurities in the H<sub>2</sub>O<sub>2</sub> product.

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Identical conditions were used for each of the resins and the analysis of H<sub>2</sub>O<sub>2</sub> was made before and after the purification column

Various types of resins were investigated for their suitability as exchange resins in  $H_2O_2$  ultrapurification processes. The resins investigated are set forth below:

#### Cationic resin (nuclear grade)

- DOWEX C75 NGH (Polystyrene-DVB macroporous 20% DVB, sulfonic, 1.7 eq H+/ℓ.)
- DOWEX MONOSPHERE C650 UPN (polystyrene-DVB gel, sulfonic, 1.9 eq H+/ℓ.)

#### Anionic resin

- DOWEX MONOSPHERE A550 UPN (polystyrene-DVB gel, quaternary ammonium, 1.0 eq OH¹/ℓ.) nuclear grade.
- AMBERLITE IRA 958 de Rohm & Haas (polyacrylic-DVB macroporous, quaternary ammonium 0.8 eq Cℓ⁻/ℓ.)

#### Chelate resin

- CHELEX 100 from Bio-Rad, amino-diacetic, 0.4 eq Na⁺/ℓ.

#### Mix bed resin

50/50 DOWEX MONOSPHERE A550 and C650.

750 m/ of  $H_2O_2$  was purified for each resin type in an ion exchange column. Three 125 m/ samples were taken for analysis, one sample from each 250 m/.

Analysis was carried out using inductively coupled plasma (ICP)/mass spectroscopy (MS) by direct introduction of  $H_2O_2$  samples. The following table summarizes the results obtained for each of the resins. The values obtained for the  $H_2O_2$  HP grade used at the entrance of the purification column are also provided. The results are shown in the Table below.

ppt(ng/kg)			CATIONIC		ANIONIC		MIXED BED	H <sub>2</sub> O <sub>2</sub> (HP grade)	CHELEX 100
Mass	Elem	H <sub>2</sub> O <sub>2</sub> (HP grade)	C75NGH	C650	IRA958	A550	C650/A550		
23	Na	34230	66	99	10363	34857	152	37040	42660
24	Mg	33	3.3	6.9	193	1073	15	34	2.4
27	Al	2096	2516	2381	237	354	218	2774	2554
39	К	68	18 ,	29	125	83	20	67	4602
40	Ca	84	13	17	164	949	19	121	27

(continued)

ppt(ng/kg)		CATIONIC		ANIONIC		MIXED BED	H <sub>2</sub> O <sub>2</sub> (HP grade)	CHELEX 100	
Mass	Elem	H <sub>2</sub> O <sub>2</sub> (HP grade)	C75NGH	C650	IRA958	A550	C650/A550		
52	Cr	47	37	45	10	13	14	55	49
55	Mn	6.6	3.1	2.7	20	11	4.4	9.4	3.8
56	Fe	209	278	404	74	271	128	261	292
58	Ni	50	15	29	59	66	16	53	11
59	Со	4.1	2.4	3.4	6.1	21	2.6		
63	Cu	7.7	6.6	16	7.0	6.8	6.3	8.9	6.1
66	Zn	506	3.7	3.2	535	500	3.9		

For the single columns investigated, the best results were obtained for the mixed bed column. With the mixed bed,

20 impurity concentration values of less than 200 ppt were obtained for each element except At with one purification of HP grade H<sub>2</sub>O<sub>2</sub> (<100 ppb).

The cationic resins were able to purify the  $H_2O_2$  except for  $A\ell$  and Fe for which an increase was observed. (These two elements are not retained on cationic resins.)

With the anionic resins. At and Fe were purified, as expected. However, the level obtained was still high (>200 ppt) Moreover an increase of Mg. K. and Ca was observed which probably stems from the chemical preparation of the resins prior to  $H_2O_2$  processing. Poor results were also observed with chelate resin.

The present invention also contemplates a system and method for the on-site preparation of ultra-high-purity mixed cleanup solutions. This aspect of the invention involves the combination of on-site ultrapurified hydrogen peroxide with an acid or base which has also been ultrapurified on-site. The described system and method find particular applicability when used in a semiconductor manufacturing facility.

Of the mixed cleanup solutions which can be prepared in accordance with the invention, preparation of the RCA acidic cleanup and the RCA basic cleanup on-site at a semiconductor fabrication facility is of particular interest. The chemicals involved in the RCA cleanup process include the following: (1) a solvent wash, such as tetrachloroethylene or a comparable solvent, to remove gross organic contaminants; (2) basic cleanup including treatment with NH<sub>4</sub>OH + H<sub>2</sub>O<sub>2</sub> + H<sub>2</sub>O; and (3) an acid cleanup with HCℓ + H<sub>2</sub>O<sub>2</sub> + H<sub>2</sub>O). See, W. Runyan et al., Semiconductor Integrated Circuit Processing Technology (1990), which is hereby incorporated by reference. For semiconductor manufacturing, such cleanup reagents are normally purchased as packaged containers. However, this implies that some handling of the solutions in those containers will be necessary, both at the manufacturer's plant and at the place of use. As noted above, such handling of ultra-high purity chemicals is undesirable.

Various other cleanup chemistries have also been proposed. For example, the Shiraki cleanup is an aggressive, pre-epitaxy cleanup. This process adds a nitric acid treatment step to the cleanup sequence, and uses somewhat higher temperatures and concentrations than the RCA cleanup process. See Ishizaki and Shiraki, Low Temperature Surface Cleaning of Silicon and its Application to Silicon MBE, 133 J. ELECTROCHEM. Soc. 666 (1986), which is hereby incorporated by reference.

The RCA basic cleanup solution typically consists of  $NH_4OH + H_2O_2 + H_2O$  in proportions of 1:1:5 or 1:2:7. In accordance with the invention, the RCA basic cleanup (or analogous cleanup solutions) can be generated on-site. Such solutions are formed by the combination of ultra-pure ammonia with ultra-pure hydrogen peroxide, both of which have been purified on-site. By the present invention, chemical purity is increased, while the risk of undetected accidental contamination is reduced.

The RCA acid cleanup solution typically includes  $HC\ell + H_2O_2 + H_2O$  in proportions of 1:1:6 or 1:2:8. The RCA acid cleanup (or analogous cleanup solutions) can be generated on-site at a wafer manufacturing plant, by combination of ultra-pure  $HC\ell$  with ultra-pure hydrogen peroxide, both of which have been purified on-site. Thus purity is increased, and the risk of undetected accidental contamination is reduced.

FIG. 2 illustrates the generation of an RCA cleanup solution on-site, at a wafer fabrication facility, using two components (30%  $\rm H_2O_2$  and 29%  $\rm NH_4OH_{aq}$  or 37%  $\rm HC\ell_{aq}$ ) in addition to ultrapure water. The two components are preferably ultrapurified on-site prior to mixing in the cleaning station.

FIG. 3 shows a wafer cleaning system in a semiconductor fabrication line. According to the illustrated embodiment, wafer or wafer batch 51 is held on a wafer support 52 and conveyed from one work station to the next by a robot 63

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or some other conventional means for achieving sequential treatment. The means of conveyance may be totally automated, partially automated, or not automated at all.

The first unit in the cleaning line is a photoresist stripping station 41, in which aqueous hydrogen peroxide 42 and sulfuric acid 43 are combined and applied to the semiconductor surface to strip off the resist. Next in line is a rinse station 44, where deionized water is applied to rinse off the previously applied stripping solution. Immediately downstream of the rinse station 44, is a cleaning station 45. In cleaning station 45, an aqueous solution of ammonia and hydrogen peroxide are applied to a wafer or wafer batch 51.

The ammonia/hydrogen peroxide solution can be supplied in two ways. According to a first method, aqueous ammonia 31 from a dissolving unit is combined with aqueous hydrogen peroxide 46, and the resulting mixture 47 is directed to cleaning station 45. In the second method, pure gaseous ammonia 32 is bubbled into an aqueous hydrogen peroxide solution 48 to produce a mixture 49, which is directed to cleaning station 45.

Once cleaned with the ammonia/hydrogen peroxide combination, the semiconductor passes to a second rinse station 50 where deionized water is applied to remove the cleaning solution. The next station is an additional cleaning station 54, in which aqueous solutions of HC $\ell$  55 and H $_2$ O $_2$  56 are combined and applied to the semiconductor surface for further cleaning.

The purified HC $\ell$  for the acid cleaning station 54 may be prepared and supplied on-site in a manner similar to that of the hydrogen peroxide purification system of FIG. 1. This step is followed by a final rinse station 57 where deionized water is applied to remove the HC $\ell$  and H $_2$ O $_2$ , and finally a drying station 58, in which the wafer or wafer batch 51 is dried.

The system shown in FIG. 3 is exemplary of cleaning lines used in semiconductor fabrication. In practice, cleaning lines can vary widely from the specific system shown in FIG. 3. For example, one or more of the units shown in FIG. 3 can be eliminated. Conversely, units not shown in FIG. 3 can optionally be added. While the specific configuration of the processing tool may vary, the system and method for on-site preparation of high-purity hydrogen peroxide and reagents including hydrogen peroxide in accordance with the invention are applicable to all such systems.

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The use of ammonia and hydrogen peroxide as a semiconductor cleaning medium at workstations such as cleaning station 45 in FIG. 3 is well known. While the proportions of the cleaning medium components vary, an exemplary system uses deionized water, 29 wt% ammonium hydroxide and 30 wt% hydrogen peroxide, combined in a volume ratio of 6: 1:1. This cleaning agent is used to remove organic residues. When used in conjunction with ultrasonic agitation at frequencies of approximately 1 MHz, particles down to the submicron range can be removed.

The on-site system for ultrapurification of  $H_2O_2$  and generation of ultrapure cleaning solutions can be connected to a point of use in the production line by piping which prevents exposure of the chemicals to an uncontrolled ambient and does not introduce contamination. In most applications, stainless steel or polymers such as high density polyethylene or fluorinated polymers are suitable materials of construction for the transfer lines.

The distance of travel between the unit and the production line can be short (e.g., in the case of a dedicated point-of-use mixing facility). However, in a more preferable embodiment, the ultrapure cleaning solution generator can be connected to multiple points of use. The generation system and points of use can be connected through ultraclean piping.

In large systems, intermediate holding tanks can be used to average the flow rate in order to compensate for varying demand. In any case, the cleaning solutions are maintained in an ultrapure environment, and are not exposed to ambient contamination. As a result, the risks of contamination due to packaging, transport, and transfer between containers can be avoided.

The distance between the point at which a cleanup solution leaves the generation system and the point of use in the production line can vary, for example, from one foot (30 cm) up to 1,000 meters or more. Greater distances are typically used when ultraclean piping is routed between buildings at a single manufacturing site.

Due to the proximity of the ultrapure purification, generation and/or mixing units to the production line, deionized water (purified in accordance with well known semiconductor manufacturing standards) is readily available for purposes such as concentration adjustment, flushing, and dissolution of gases.

The standards commonly used in the semiconductor manufacturing industry are well known among those skilled in the art. For example, typical standards for the purity of water used in these semiconductor manufacturing processes include a resistivity of at least about 15 megohm-cm at 25°C (typically 18 megohm-cm at 25°C), less than about 25 ppb of electrolytes, a particulate content of less than about 150/cm³, a particle size of less than 0.2 micron, a microorganism content of less than about 10/cm³, and total organic carbon of less than 100 ppb. Additives can be introduced into the purification water if desired.

In the method and system according to the invention, a high degree of control over the product concentration and, hence, flow rates is preferably maintained, by precise monitoring and metering using known equipment and instrumentation. A convenient means of achieving such control is ultrasonic wave propagation to monitor density. Other methods will be readily apparent to those skilled in the art.

In summary, the invention can be used as an on-site system for providing ultrapure chemicals to a point of use. The purification system can also be adapted to operate as part of a manufacturing unit for producing ultra-high-purity

chemicals which are to be shipped. However, when used in this manner, many of the advantages gained with on-site purification cannot be realized. This is a result of the inherent risks in the handling of ultra-high-purity chemicals. Nevertheless, for customers requiring packaged chemicals, the inventive system and method at least provide an initial chemical purity which is higher than that available by other preparation techniques. Again, in such applications a dryer stage may also be used after the ionic purifier.

While the invention has been described in detail with reference to specific embodiments thereof, it will be apparent to those skilled in the art that various changes and modifications can be made, and equivalents employed, without departing from the scope of the appended claims. For example, the disclosed innovative techniques can be applied to the manufacture of products other than ICs, such as discrete semiconductor components (e.g., optoelectronic and power devices), and to other manufacturing technologies in which IC manufacturing methods have been adopted (e.g., the manufacture of thin-film magnetic heads and active-matrix liquid-crystal displays).

#### Claims

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- 1. An on-site subsystem for providing ultra-high-purity reagents comprising H<sub>2</sub>O<sub>2</sub> to a point of use, comprising:
  - (a) a tank connected to receive aqueous H2O2 and to provide a flow of H2O2 therefrom;
  - (b) an anionic exchange bed and a cationic exchange bed connected to receive said flow of  $H_2O_2$  from said tank, and to produce therefrom a purified  $H_2O_2$  flow with a reduced level of ionic contaminants, wherein said cationic exchange bed is preconditioned with acid and said anionic bed is preconditioned with bicarbonate ions;
    - (c) a filter downstream of said anionic exchange bed and said cationic exchange bed; and
    - (d) piping which routes said aqueous H<sub>2</sub>O<sub>2</sub> from said filter to said point of use.
- 25 2. The one-site subsystem according to claim 1, further comprising one or more additional anionic exchange beds and one or more additional cationic exchange beds.
  - 3. The one-site subsystem aaccording to claim 1 or 2, wherein the subsustem and the point of use are located in a semiconductor fabrication facility.
  - 4. An one-site subsystem for providing ultra-high-purity reagents comprising H<sub>2</sub>O<sub>2</sub> to a point of use, comprising:
    - (a) a tank connected to receive aqueous  $\rm H_2O_2$  and to provide a flow of  $\rm H_2O_2$  therefrom:
    - (b) a mixed anionic/cationic exchange bed connected to receive said flow of  $H_2O_2$  from said tank, and to produce therefrom a purified  $H_2O_2$  flow with a reduced level of ionic contaminants:
    - (c) a filter downstream of said anionic/cationic mixed exchange bed; and
    - (d) piping which routes said aqueous H<sub>2</sub>O<sub>2</sub> from said filter to said point of use.
- 5. The on-site subsystem according to claim 4, further comprising one or more anionic exchange beds and one or more cationic exchange beds.
  - 6. The one-site subsystem according to claim 4 or 5, wherein the subsystem and the point of use are located in a semiconductor fabrication facility.
- 7. An one-site subsystem for providing ultra-high-purity reagents comprising H<sub>2</sub>O<sub>2</sub> to a point of use, comprising:
  - (a) a tank connected to receive aqueous  $\rm H_2O_2$  and to provide a flow of  $\rm H_2O_2$  therefrom;
  - (b) an anionic exchange bed and a cationic exchange bed connected to receive said flow of  $H_2O_2$  from said tank, and to produce therefrom a purified  $H_2O_2$  with a reduced level of ionic contaminants;
  - (c) an ionic purifier system, connected to pass a gaseous reagent precursor through an area of gas/liquid contact to produce an ultrapure gaseous reagent;
  - (d) generation an dmixing subsystems, connected to combine said ultrapure gaseous reagent with deionozed water and with said purified  $H_2O_2$  flow to produce an ultrapure cleanup solution; and
  - (e) piping which routes said ultrapure cleanup solution to the point of use.
  - 8. The one-site subsystem according to claim 7, further comprising one or more additional anionic exchange beds and one or more additional cationic exchange beds.

- 9. The one-site subsystem according to claim 7 or 8, wherein the subsystem and the point of use are located in a semiconductor device fabrication facility.
- 10. The one-site subsystem according to one of claims 7 to 9, wherein the generation and mixing subsystems are separate.
- 11. The one-site subsystem according to one of claims 7 to 10, wherein the generation and mixing subsystems are combined.
- 10 12. The one-site subsystem according to one of claims 7 to 11, wherein the gaseous reagent is Hcl.
  - 13. The one-site subsystem according to one of claims 7 to 12, wherein the gaseous reagent is NH<sub>3</sub>.
- 14. A method for the one-site provision of ultra-high-purity reagents comprising H<sub>2</sub>O<sub>2</sub> to a point of use, comprising the steps of:
  - (a) providing a flow of aqueous H<sub>2</sub>O<sub>2</sub> from a tank which is located at the same site as the point of use;
  - (b) passing said flow of  $H_2O_2$  through an anionic exchange bed and a cationic exchange bed, to produce a purified  $H_2O_2$  flow with a reduced level of ionic contaminants; wherein said cationic exchange bed is precon-
  - ditioned with acid and said anionic bed is preconditioned with bicarbonate ions;

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- (c) filtering said purified flow, to produce a flow of ultra-purity aqueous H<sub>2</sub>O<sub>2</sub> solution; and
- (d) routing said flow of ultra-purity aqueous H<sub>2</sub>O<sub>2</sub> solution through piping to the point of use.
- 15. The method according to claim 14, further comprising passing said flow of H<sub>2</sub>O<sub>2</sub> through one or more additional anionic exchange beds and through one or more additional cationic exchange beds.
  - **16.** The method according to claim 14 or 15, wherein the point of use is located in a semiconductor device fabrication facility.
- 39 17. A method for the one-site provision of ultra-high-purity reagents comprising H<sub>2</sub>O<sub>2</sub> to a point of use, comprising the steps of:
  - (a) providing a flow of aqueous  $H_2O_2$  from a tank which is located at the same site as the semiconductor manufacturing operation:
  - (b) passing said flow of  $H_2O_2$  through a mixed anionic/cationic exchange bed, to produce a purified  $H_2O_2$  flow with a reduced level of ionic contaminants:
  - (c) filtering said purified flow, to produce a flow of ultra-purity aqueous H<sub>2</sub>O<sub>2</sub> solution; and
  - (d) routing said flow of ultra-purity aqueous H<sub>2</sub>O<sub>2</sub> solution through piping from said filter to the point of use.
- 40 18. The method according to claim 17, further comprising passing said flow of H<sub>2</sub>O<sub>2</sub> through one or more anionic exchange beds and through one or more cationic exchange beds.
  - 19. The method according to claim 17 or 18,, wherein the point of use is located in a semiconductor device fabrication facility.
  - 20. A method for the one-site provision of ultra-high-purity reagents comprising H<sub>2</sub>O<sub>2</sub> to a point of use, comprising the steps of:
    - (a) providing a flow of aqueous H<sub>2</sub>O<sub>2</sub> from a tank which is located at the same site as the point of use;
    - (b) passing said flow of H<sub>2</sub>O<sub>2</sub> through an ionic exchange bed and a cationic exchange bed, to produce a purified H<sub>2</sub>O<sub>2</sub> flow with a reduced level of ionic contaminants, wherein said cationic exchange bed is preconditioned with bicarbonate ions:
      - (c) passing a gaseous reagent precursor through an area of gas/liquid contact in an ionic purifier system to produce an ultrapure gaseous regent;
      - (d) combining said ultrapure gaseous reagent with deionized water and with said purified  $H_2O_2$  flow to produce an ultrapure cleanup solution; and
      - (e) routing said flow of ultra-purity cleanup solution to the point of use.

- 21. The method according to claim 20, further comprising passing said flow of H<sub>2</sub>O<sub>2</sub> through one or more additional anionic exchange beds and through one or more additional cationic exchange beds.
- 22. The method according to claim 20, wherein the point of use is located in a semiconductor device fabrication facility.
- 23. The method according to claim 21 or 22, wherein the gaseous reagent is Hcl.
- 24. The method according to one of claims 21 to 23, wherein the gaseous reagent is NH<sub>3</sub>
- 25. A method for preparing ultra-high-purity reagents comprising H<sub>2</sub>O<sub>2</sub>, comprising the steps of:
  - (a) providing a flow of aqueous H<sub>2</sub>O<sub>2</sub> from a tank;
  - (b) passing said flow of  $H_2O_2$  through an anionic exchange bed and a cationic exchange bed, to produce a purified  $H_2O_2$  flow with a reduced level of ionic contaminants; wherein said cationic exchange bed is preconditioned with acid and said anionic bed is preconditioned with bicarbonate ions; and
  - (c) filtering said puritied flow, to produce a flow of ultra-purity aqueous H<sub>2</sub>O<sub>2</sub> solution.
  - 26. A method for preparing ultra-high-purity reagents comprising H<sub>2</sub>O<sub>2</sub>, comprising the steps of:
- 20 (a) providing a flow of aqueous H<sub>2</sub>O<sub>2</sub> from a tank;
  - (b) passing said flow of  $H_2O_2$  through an anionic exchange bed and a cationic exchange bed, to produce a purified  $H_2O_2$  flow with a reduced level of ionic contaminants; wherein said cationic exchange bed is preconditioned with acid and said anionic bed is preconditioned with bicarbonate ions:
  - (c) passing a gaseous reagent precursor through an area of gas/liquid contact in an ionic purifier system to produce an ultrapure gaseous reagent; and
  - (d) combining said ultrapure gaseous reagent with deionized water and with said purified H<sub>2</sub>O<sub>2</sub> flow to produce an ultrapure cleanup solution.

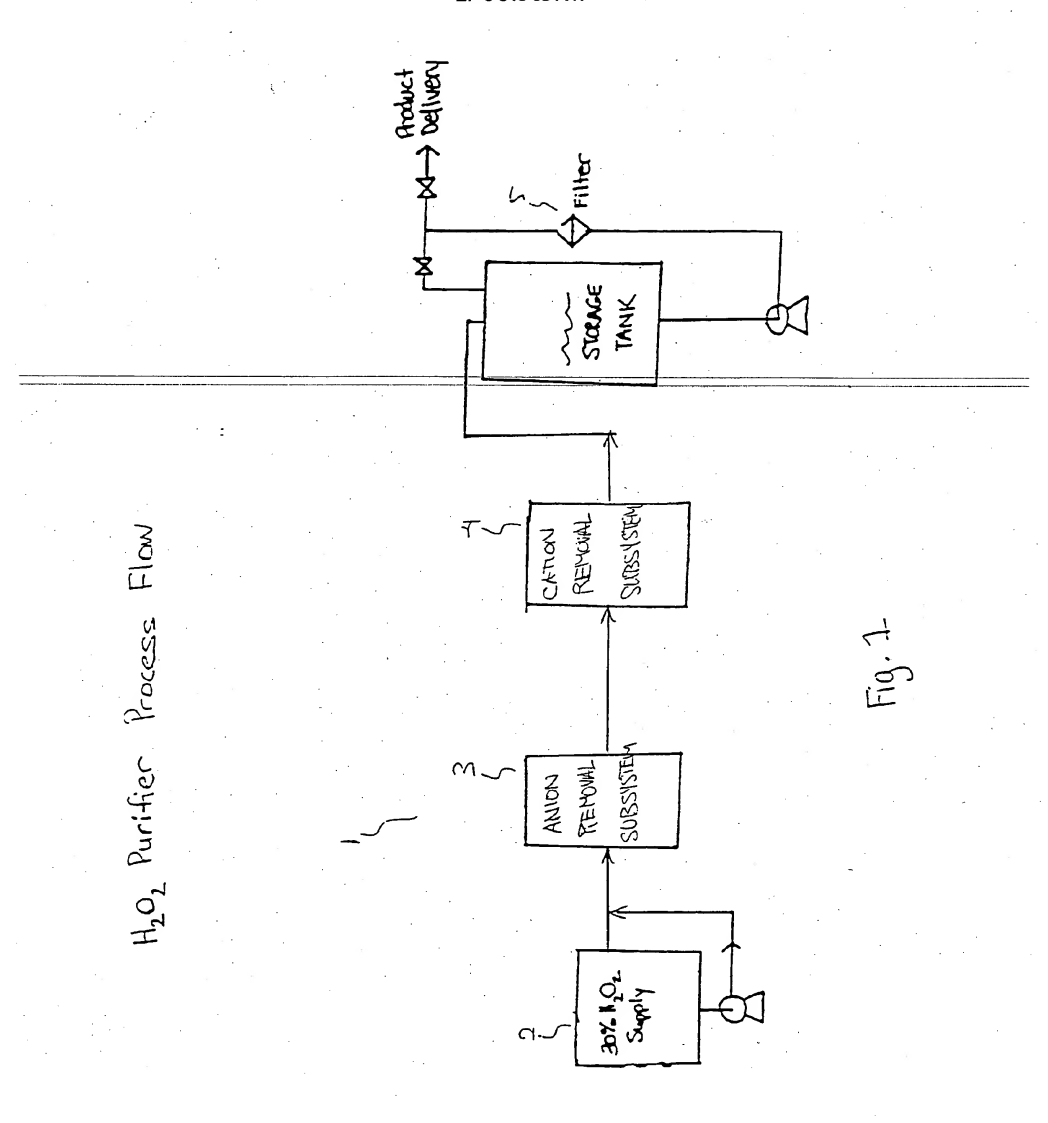
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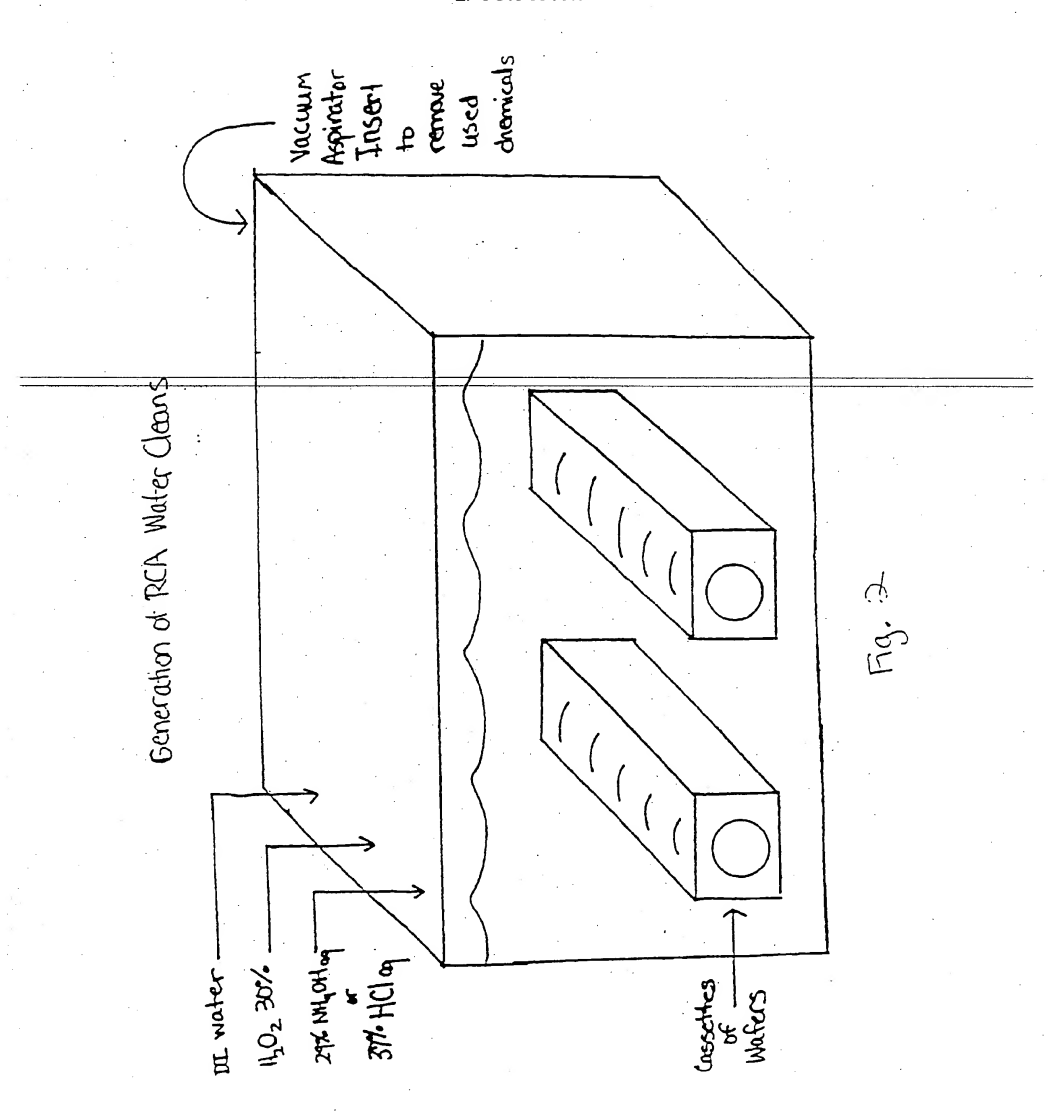
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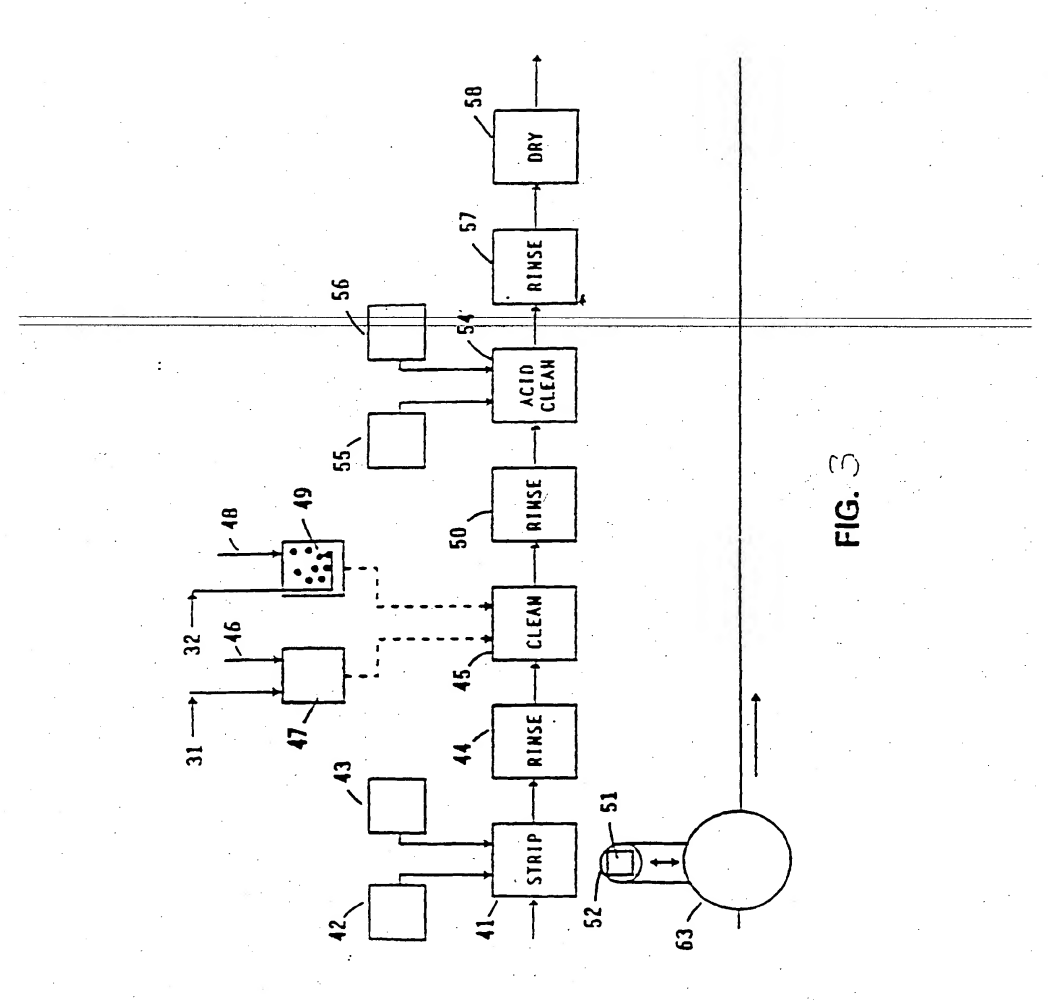
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## EUROPEAN SEARCH REPORT

Application Number

EP 97 40 2937

ategory	Citation of document with indication, of relevant passages	where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.6)
Ρ, χ	WO 96 39237 A (STARTEC VE ;HOFFMAN JOE G (US); CLAF December 1996	1,3,7,	C01B15/013	
·, γ	* claims 1-7 * * claims 1-7 *		2,4-6,8, 15,18,21	
1	DATABASE WPI		4-6	
	Section Ch, Week 9648 Derwent Publications Ltd Class E36, AN 96-481970 XP002057184 & JP 08 245 204 A (MITSUE INC), 24 September 1996 * abstract *	BISHI GAS CHEM CO		
1	WO 92 06918 A (INTEROX CI April 1992 * page 7, line 19 - page	2.5,8, 15,18,21		
<b>A</b>	DATABASE WPI Section Ch, Week 9646 Derwent Publications Ltd Class A91, AN 96-461142 XP002057185 & JP 08 231 207 A (MITSUITO September 1996 * abstract *	., Landon, GB;	4,6	TECHNICAL FIELDS SEARCHED (Int.Cl.6)
A	WO 91 01267 A (INTEROX C February 1991 * page 7, line 8 - line		2,5,8, 15,18,21	
	The present search report has been dra	wn up for all claims		<u>.</u>
	Place of search	Date of completion of the search		Examiner
	THE HAGUE	27 February 1998	Ros	is, R
X : par Y : par doc	CATEGORY OF CITED DOCUMENTS  tocutarly relevant if taken alone ticularly relevant if combined with another turnent of the same category hnological background	T : theory or principle E earlier patent doc after the tiling date D : document cited in L : document cited fo	:ument, but puble e n the application or other reasons	lished on, or